

## **Product Description**

Pioneering GTPase and Oncogene Product Development since 2010

### CDC42(G12V) PROTEIN

### Cdc42(G12V) Mutant

Cat. #: 10109

Product Name: Cdc42 Protein G12V mutant

Synonyms: Cell division cycle 42, G25K, CDC42Hs

Source: Human, recombinant full length, His6-tag

Expression Host Species: E. col

Molecular Weight: 21 kDa

Purity: >95% by SDS-PAGE

**Introduction:** Small GTPases are a super-family of cellular signaling regulators. Cdc42 belongs to the Rho sub-family of GTPases that regulate cell motility, cell division, and gene transcription. GTP binding increases the activity of Cdc42, and the hydrolysis of GTP to GDP renders it inactive. GTP hydrolysis is aided by GTPase activating proteins (GAPs), while exchange of GDP for GTP is facilitated by guanine nucleotide exchange factors (GEFs).

Amino Acid Sequence (1-191, G12V)

MQTIKCVVVGDVAVGKTCLLISYTTNKFPSEYVPTVFDNYAVTVMIGGEPYTLGLFDTAGQEDYDRL RPLSYPQTDVFLVCFSVVSPSSFENVKEKWVPEITHHCPKTPFLLVGTQIDLRDDPSTIEKLAKNKQ KPITPETAEKLARDLKAVKYVECSALTQKGLKNVFDEAILAALEPPEPKKSRRCVLL

#### **Properties**

Physical Appearance (form): Dissolved in 20mM Tris-HCl, pH8.0, 150mM NaCl.

Physical Appearance (form): White or clear

**Concentration:** 1 mg/mL

#### Storage: -80°C

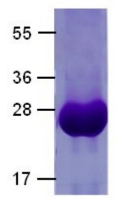
#### **Preparation Instructions:**

Centrifuge the vial before open the cap and reconstitute in water. Adding of 10 mM βmercaptoethanol or 1 mM DTT into the solution to protect the protein is recommended and using of non-ionic detergents such as n-Dodecyl β-D-maltoside (DoDM) or polyethylene detergents (e.g. C12E10) also help to stabilize the protein. Avoid repeated freezing and thawing after reconstitution. The purity of His-tagged Cdc42 G12V was determined by SDS-PAGE and Coomassie Brilliant Blue Staining.



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#### **References:**

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- 3. Kawasaki, Y. et al., Oncogene 26: 7620-7627, 2007.
- 4. Manser, E. et al., Nature 363: 364-367, 1993.
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- 7. Shen, Y. et al., Dev. Cell 14: 342-353, 2008.
- 8. Wu, W. J. et al., Nature 405: 800-804, 2000.
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